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L1
             48 S PHOTOLYSIS AND METAL ALKYL
L2
              0 S L2 AND CARBOXYLIC ACID
L3
              0 S L2 AND ORGANIC ACID
L4
L5
              0 S L2 AND FORMIC ACID
              0 S L2 AND ACETIC ACID
1.6
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       1644131 METAL
        567045 ALKYL
          1739 METAL ALKYL
                 (METAL (W) ALKYL)
        352091 ORGANIC
       4142789 ACID
          3358 ORGANIC ACID
                 (ORGANIC (W) ACID)
L7
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                 (METAL (W) ALKYL)
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L8
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D1-20 IS NOT A RECOGNIZED COMMAND
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For a list of commands available to you in the current file, enter
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     ANSWER 1 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
L8
     2004:955757 CAPLUS
AN
DN
     142:241258
TΤ
     Polyamide resin composition
     Jung, Yong Gyun; Lee, Do Geun; Park, Eun Ha
TN
PΑ
     Kolon Ind. Inc., S. Korea
SO
     Repub. Korean Kongkae Taeho Kongbo, No pp. given
     CODEN: KRXXA7
DT
     Patent
LA .
     Korean
FAN.CNT 1
                         KIND DATE
                                          APPLICATION NO.
                                                                   DATE
     PATENT NO.
                                            ------
                                                                   _____
                         ----
     KR 2003053835
                         Α
                                20030702 KR 2001-83850
                                                                   20011224
PΤ
PRAI KR 2001-83850
                                20011224
     A polyamide resin composition is provided, which shows excellent mech. strength
     and surface characteristics and exhibits an excellent coating property by
     only base coating without a primer treatment. The polyamide resin composition
     comprises 60-89.5% a polyamide; 5-15% a thermoplastic rubber; 5-20% a
     copolymer of an aromatic vinyl compound and Ph maleimide; and 0.5-5% an alkali
     metal alkyl sulfate. Preferably the polyamide is
     polyamide 66 or polyamide 6; the thermoplastic elastomer is an
     ethylene-propylene-diene copolymer grafted with 0.4-2%
     \alpha, \beta-unsatd. carboxylic acid, anhydride or
     their derivs.; the copolymer of an aromatic vinyl compound and Ph maleimide is
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poly(Ph maleimide styrene) with number average mol. weight 80000-200000, which is

grafted with 40-60 parts Ph maleimide, 40-60 parts styrene, and 1-5 parts

maleic anhydride; and the alkali metal alkyl sulfate

compound is a sodium C5-C25 alkyl sulfate.

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ANSWER 2 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
L8 ·
AN
     1999:819244 CAPLUS
DN
     132:49829
     Preparation and formulation of naphthalenesulfamylmethyl-carbapenems for
TI
     use as antibacterial agents
     Ratcliffe, Ronald W.; Wilkening, Robert R.; Blizzard, Timothy A.
IN
PΑ
     Merck & Co., Inc., USA
SO
     PCT Int. Appl., 60 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
FAN.CNT 1
                                DATE
                                             APPLICATION NO.
                                                                    DATE
     PATENT NO.
                         KIND
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     WO 9966928
                                19991229
PΙ
                          A1
                                             WO 1999-US14223
                                                                    19990623
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             GE, HR, HU, ID, IL, IN, IS, JP, KG, KR, KZ, LC, LK, LR, LT, LV,
             MD, MG, MK, MN, MX, NO, NZ, PL, RO, RU, SG, SI, SK, TJ, TM, TR,
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TT, UA, US, UZ, VN, YU, ZA, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG CA 1999-2335507 CA 2335507 AA19991229 19990623 AU. 9947110 Α1 20000110 AU 1999-47110 19990623 20010411 EP 1089730 A1 EP 1999-930606. 19990623 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE, SI, LT, LV, FI, RO US 6294528 20010925 US 1999-338644 19990623 B1JP 2002518447 **T2** 20020625 JP 2000-555614 19990623 Р 19980625

PRAI US 1998-90684P W WO 1999-US14223 19990623

OS MARPAT 132:49829

GI

AB Naphthalenesulfamylmethyl-carbapenems I [R = H, aryl, heterocyclyl, alkenyl, etc.; R1 = H, Me; R2 = H, OH, F, protected hydroxy; M = H, alkali metal, alkyl, alkenyl, etc.;] were prepared for use as antibacterial agents. Thus, I [R = H, R1 = Me, R2 = OH, M = Na] was prepared starting from (4S,5R,6S)-3-(hydroxymethyl)-4-methyl-7-oxo-6-[(1R)-1-[[(2-propenyloxy)carbonyl]oxy]ethyl]-1-azabicyclo[3.2.0]hept-2-ene-2carboxylic acid 2-propenyl ester and Pharmaceutical formulations of 6H-dibenzo[c,e][1,2]thiazine, 5,5-dioxide. the prepared carbapenems were also presented.

RE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

I

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L8
    ANSWER 3 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
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ΑN 1995:761984 CAPLUS

DN 123:287187

Preparation of fluorenyl-containing metallocenes and their use in TI polymerization of olefins

IN Alt, Helmut G.; Palackal, Syriac J.; Patsidis, Konstantinos; Welch, M. Bruce; Geerts, Rolf L.; Hsieh, Eric T.; McDaniel, Max P.; Hawley, Gil R.; Smith, Paul D.

Phillips Petroleum Co., USA
U.S., 14 pp. Cont.-in-part of U.S. 5,191,132.
CODEN: USXXAM so

 \mathtt{DT} Patent LA English FAN.CNT 21

PΑ

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US 1998-85945
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                            19980528
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OS MARPAT 123:287187

A fluorenyl precursor is treated with an alkali metal alkyl in a hydrocarbon or noncyclic ether diluent and the resulting fluorenyl salt is treated with a Group IVB, VB, or VIB metal compound to give a metallocene. Thus, fluorene was treated successively with BuLi and BrCH2CH2Br to give 1,2-di-9-fluorenylethane, which was treated with MeLi in toluene or Et2O and then with ZrCl4 to give a bridged zirconocene. This metallocene was used with methylaluminoxane to polymerize ethylene at 90° to give a polyethylene with weight-average mol. weight 633 + 103 and d. 0.9384.

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L8 ANSWER 4 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
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AN 1995:699539 CAPLUS

DN 123:255879

AΒ

TI Mechanistic aspects of Ru(BINAP)-catalyzed asymmetric hydrogenation of vinyl carboxylic acid derivatives

AU Chan, Albert S. C.; Chen, Chih Chiang; Yang, Teng Kuei; Huang, Jen Hwei; Lin, Ying Chih

CS Department of Applied Biology and Chemical Technology, Hong Kong Polytechnic University, Hong Kong, Hong Kong

SO Inorganica Chimica Acta (1995), 234(1-2), 95-100 CODEN: ICHAA3; ISSN: 0020-1693

PB Elsevier

DT Journal

LA English

OS CASREACT 123:255879

ABThe mechanism of the RuII(BINAP)-catalyzed hydrogenation of vinyl carboxylic acids has been investigated via detailed deuterium labeling studies. The activation of H2 by the Ru catalyst was found to be via a heterolytic splitting route. The regioselectivity of the hydride migration step strongly correlates to the stability of the resulting metal-alkyl intermediate. When an α, β unsatd. carboxylic acid such as 2-(6-methoxy-2naphthyl)acrylic acid is used as the substrate, the hydride migrates exclusively to the α position, forming a metal-primary alkyl intermediate. In the case of a β, γ -unsatd. carboxylic acid with an electron-withdrawing group attached to the β -carbon, the hydride migrates exclusively to the γ -carbon, forming a five-membered ring metal-alkyl intermediate with the β -carbon coordinated to the metal. The product formation step involves two competing routes: the hydrogenolysis and the solvolysis of the metal-alkyl intermediates. The choice of each route is highly dependent on the reaction conditions. The solvolysis route is significant if the reaction is carried out under low H2 pressure and the reaction medium is more acidic. Under high H2 pressure, hydrogenolysis becomes the dominant route. Under basic conditions, the solvolysis route is essentially shut off and only the hydrogenolysis product is obtained. A unified mechanism which explains all of the known

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L8 ANSWER 5 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
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AN 1993:101677 CAPLUS

DN 118:101677

TI Preparation of fluorene derivatives .

exptl. results is proposed.

IN Patsidis, Konstantinos; Palackal, Syriac Joseph; Alt, Helmut

PA Phillips Petroleum Co., USA

SO Eur. Pat. Appl., 17 pp.

CODEN: EPXXDW

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                               19921110
                        A2
                                           JP 1992-116375
                                                                  19920508
    JP 05148166
                               19930615
    AT 158269
                        E
                               19971015
                                           AT 1992-107781
                                                                  19920508
                        Т3
                                           ES 1992-107781
                                                                 19920508
    ES 2106795
                               19971116
                                           KR 1992-7931
                                                                  19920508
    KR 178279
                        B1
                               19990515
PRAI US 1991-697363
                         Α
                               19910509
    MARPAT 118:101677
OS
    Title compds. ZRZ (at least 1 Z = organic radical containing cyclopentadienyl
AB
     functionality and the other Z = cyclopentadienyl-containing organic radical, Cl,
     Br, iodo) were prepared by reaction of the precursor to Z with an alkali
    metal alkyl to produce the Z anion and reaction of the
     latter with XRX (each X = Br, Cl, iodo; R = C1-20 alkylene which may
     contain Ge, B, Si, P, N, Al, O or R = Ge, Si, B, Al, P, Sn). Thus, BuLi
     was added to a solution of fluorene in THF and the mixture was stirred 1 h,
     then added over 2 h to a stirred solution of 1,2-dibromoethane in pentane.
     Work up gave 1-bromo-2-(fluorenyl)ethane.
    ANSWER 6 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
L8
    1991:559451 CAPLUS
AN
DN
    115:159451
    Process for preparing transition metal cyclopentadienyl carbonyl compounds
TТ
    Bell, Donald R.; Berris, Bruce C.
TN
     Ethyl Corp., USA
PA
    U.S., 4 pp.
SO
     CODEN: USXXAM
DT
     Patent
    English
LA
FAN.CNT 1
    PATENT NO.
                        KIND
                             DATE
                                           APPLICATION NO.
                                                                  DATE
                        ----
PΙ
    US 5026885
                        Α
                               19910625
                                           US 1990-488886
                                                                  19900306
                        AA
                                           CA 1991-2036861
                                                                  19910220
    CA 2036861
                               19910907
    CA 2036861
                        C
                               20001024
    JP 04211693
                        A2
                               19920803
                                           JP 1991-60975
                                                                  19910304
                               19910911
    EP 446007
                        A1
                                           EP 1991-301793
                                                                  19910305
    EP 446007
                        B1
                              19940907
        R: BE, DE, ES, FR, GB, IT
    AU 9172073
                        A1
                               19910912
                                           AU 1991-72073
                                                                  19910305
     AU 634865
                         B2
                               19930304
PRAI US 1990-488886
                               19900306
                         Α
    MARPAT 115:159451
OS
AB
    Title compds. [RxCpM(CO)y]n (R = hydrocarbyl; Cp = C5H5 =
     \eta5-cyclopentadienyl; M = transition metal, e.g., Mn, Co, Mo, Ru, Rh; x
     = 0-5; y = 1-7; n = 1,2; any 2 R groups together can form a fused ring
     with Cp) are prepared in one step by reaction of the corresponding
     transition metal salt of an organic carboxylic acid, a
     \beta-diketone, or a \beta-keto ester with 1-12 mol of a
     cyclopentadienyl compound and 0.3-10 mol of a metal alkyl
     reducing agent (preferably Et3Al) under CO pressure at 75-225°.
     E.g., Mn(OAc)2 10g, methylcyclopentadiene 13.9 g, and Et3Al 19.9 g in 50 ·
     mL PhMe and 12.9 g Et2O reacted under 800 psig CO at 175° 2 h to
    give 85% (C5H4Me)Mn(CO)3.
```

L8 ANSWER 7 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN AN 1991:23976 CAPLUS

DN 114:23976 Preparation of antihypercholesterinemic and antiarteriosclerotic ΤI 6-fluoro-3,5-dihydroxy carboxylic acids. Beck, Gerhard; Bartmann, Wilhelm; Wess, Guenther; Granzer, Ernold IN PA Hoechst A.-G., Germany SO Ger. Offen., 46 pp. CODEN: GWXXBX DT Patent LA German FAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE ------ - **-** -------19880806 PΙ DE 3826814 **A1** 19900208 DE 1988-3826814 19890728 EP 354418 A2 19900214 EP 1989-113917 EP 354418 Α3 19910731 EP 354418 В1 19940518 AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, LU, NL, SE AT 105838 Ε 19940615 AT 1989-113917 19890728 ES 2054947 Т3 19940816 ES 1989-113917 19890728 FI 8903677 Α 19900207 FI 1989-3677 19890803 FI 90546 В 19931115 FI 90546 C 19940225 DK 8903844 Α 19900207 DK 1989-3844 19890804 Α 19900207 NO 1989-3167 19890804 NO 8903167 В 19940801 NO 175637 С 19941109 NO 175637 **A1** 19900208 AU 1989-39297 19890804 AU 8939297 AU 616291 В2 19911024 A2 19900330 JP 1989-201500 19890804 JP 02091034 Α 19900425 ZA 1989-5959 19890804 ZA 8905959 Α US 1989-389809 19890804 US 5204351 19930420 PRAI DE 1988-3826814 Α 19880806 EP 1989-113917 Α 19890728 os CASREACT 114:23976; MARPAT 114:23976

GI

AB Title acid derivs. I [R1 = various (substituted) Ph, 3-pyridinyl, 5-pyrimidinyl, 4-pyridazinyl, 2- or 3-pyrrolyl, 2- or 3-thienyl, 3-furyl (5-membered rings may be benzo-fused), trisubstituted vinyl; R2 = H, alkyl, alkenyl, (halo- or alkyl)benzyl, alkali metal, (alkyl- or hydroxyalkyl)ammonium] and the corresponding lactones II

were prepared For example, 5-hydroxymethyl-2,4-bis-(4-fluorophenyl)-6-(1-methylethyl)pyrimidine was subjected to a sequence of oxidation to the aldehyde, condensation with Ph2P(0)CHFCO2Et, reduction with Dibal, and reoxidn. to give (2Z)-2-fluoro-3-[246-bis-(4-fluorophenyl)-6-(1-methylethyl)pyrimidin-5-yl]prop-2-enal. This was condensed with the dianion of (S)-AcoCHPhCPh2OH, followed by transesterification with NaOMe, condensation with the anion of AcoCMe3, and reduction with Et3B-NaBH4 to give I (R1 = Q, R2 = CMe3) (III). The IC50 values of III for inhibiting HMG-CoA reductase in vitro and cholesterol biosynthesis in cell culture were 2.9 + 10-9 and 1.9 + 10-8, resp. Prepns. of 13 I, 12 II, and numerous intermediates are described, with test data for 5 compds.

L8 ANSWER 8 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1989:574668 CAPLUS

DN 111:174668

TI Phosphinylcycloalkylcarbonyl- and phosphinylcycloalkenylcarbonyldipeptide as angiotensin converting enzyme inhibitors

IN Weller, Harold Norris, III; Gordon, Eric Michael

PA E. R. Squibb and Sons, Inc., USA

SO Ger. Offen., 16 pp. CODEN: GWXXBX

DT Patent

LA German

FAN CNT 1

FAN.	CNT I					
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
PI	DE 3831936	A1	19890406	DE 1988-3831936	19880920	
	US 4849525	Α	19890718	US 1987-98651	19870921	
	CA 1316630	A1 ·	19930420	CA 1988-575028	19880817	
	GB 2210043	· A1	19890601	GB 1988-21942	19880919	
	GB 2210043	B2	19910508	•		
	FR 2620711	A1	19890324	FR 1988-12268	19880920	
	FR 2620711	B1	19950331			
	JP 01113400	A2	19890502	JP 1988-237454	19880921	
PRAI	US 1987-98651	Α	19870921			
os	MARPAT 111:174668			,		
GI						

$$Q = \begin{pmatrix} Z \\ Q = \\$$

The title compds. [I; R = H, alkyl, cycloalkyl, aralkyl, aminoalkyl, AB hydroxyalkyl, mercaptoalkyl; R1 = H, alkyl, haloalkyl, indolylalkyl, phenylalkyl, imidazolylalkyl, carbamoylalkyl, etc.; R2 = alkyl, aralkyl, aminoalkyl; R3, R4 = H, alkali metal, alkyl, (substituted) alkylcarbonyloxyalkyl, phenylcarboyloxyalkyl; X = Q; Z = atoms to complete a (substituted) C3-10 cycloalkyl, C3-7 cycloalkenyl, C5-7 heterocyclyl ring; R5 = H, cyclohexyl, C1-4 alkoxy, (substituted) Ph, phenylalkyl, phenylalkoxy, phenylalkylthio], useful as angiotensin converting enzyme (ACE) inhibitors (no data) were prepared trans-6-[Hydroxy(4-phenylbutyl)phosphinyl]-3-cyclohexene-1carboxylic acid [preparation from Et (4phenylbutyl)phosphinate given] in THF at 125° was stirred 1 h with DCC and H-Ala-Pro-OBzl (Bzl = PhCH2). TsOH and Et3N were added. After adding TsOH and Et3N, the mixture was stirred for 18 h to give the coupling product as a separable mixture of isomers. Isomer A was hydrogenolyzed in MeOH over 20% Pd(OH)2/carbon followed by stirring with aqueous LiOH to give N-[trans-6-[hydroxy(4-phenylbutyl)phosphinyl]-3-cyclohexene-1-carbonyl]-Lalanyl-L-proline dilithium salt.

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ANSWER 9 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
L8
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1989:425842 CAPLUS AN

111:25842 DN

ΤI Novel collectors and processes for making and using same

Wang, Samuel Shan Ning; Nagaraj, D. R. IN

PΑ American Cyanamid Co., USA

Eur. Pat. Appl., 9 pp. SO

CODEN: EPXXDW

Patent DT English

LA FAN.CNT 1

		_										
	PA:	CENT	NO.			KIN	D	DATE	AP	PLICATION NO	•	DATE
		-		- 			-				-,	
ΡI	EР	3117	59			A2		19890419	EP	1988-112212		19880728
	EΡ	3117	59	•		. A3		19890823				
	EP	3117	59			B1		19920902				
		R:	DE,	FR,	GB,	IT,	NL					
	US	4871	466			Α		19891003	US	1987-108611		19871015
	BR	8805	264			Α		19900605	BR	1988-5264		19881013
	CA	1299	197			A1	•	19920421	CA	1988-579990		19881013
	ΑU	8823	793			A1		19890420	AU	1988-23793		19881014
	ΑU	6043	73			B2		19901213				
	US	4929	343			Α		19900529	US	1989-375443		19890705
	US	5237	079			Α		19930817	US	1990-478307		19900212
PRAI	US	1987	-1086	511		Α		19871015				
	US	1989	-3754	143		A3		19890705		•		,

Alkali metal hydroxamates are produced by reacting the Me or Et ester of a AB fatty acid having 6-22 C atoms with hydroxylamine salt and an alkali metal hydroxide in the presence of a water-C8-22 alc. mixture and, preferably, a non-ionic or cationic surfactant. The solution of hydroxamate thus produced can be used in the froth flotation of non-sulfide minerals such as kaolin. The process eliminates the need for hazardous and expensive recovery steps such as filtration and results in high conversions, i.e., 85-95%.

ANSWER 10 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN L8

AN1989:38631 CAPLUS

DN

Preparation and use of acylaminopropionates as surfactants, emulsifiers, ΤI wetteners, detergent components, and in production of waterproof leather

IN Dahmen, Kurt; Mertens, Richard; Stockhausen, Dolf

PΑ Chemische Fabrik Stockhausen G.m.b.H., Fed. Rep. Ger.

Ger. Offen., 9 pp.

CODEN: GWXXBX

DTPatent

LA German

FAN.	CNT 2				•
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
					
PI	DE 3717961	A1	19880505	DE 1987-3717961	19870527
	DE 3717961	C2	19940526		
	EP 265818	A2	19880504	EP 1987-115365	19871021
	EP 265818	A3	19900425		
	EP 265818	B1	19940928		
	R: AT, BE, CH,	DE, ES	, FR, GB, G	R, IT, LI, NL, SE	
	JP 63112544	A2	19880517	JP 1987-266178	19871021
	JP 2577011	B2	19970129		
	ES 2003836	Т3	19941216	ES 1987-115365	19871021
	AU 8780101	A1	19880428	AU 1987-80101	19871023
	AU 602171	B2	19901004		
	BR 8705693	Α	19880531	BR 1987-5693	19871023
	SU 1833368	A3	19930807	SU 1989-4613176	19890104
	RU 2062302	C1	19960620	RU 1989-4613251	19890104
•	LV 11044	В	19961020	LV 1993-715	19930628
	LT 3617	В	19951227	LT 1993-1535	19931206
	LT 3805	В	19960325	LT 1993-1597	19931215
PRAI	DE 1986-3636497	A1	19861027		
	DE 1987-3717961	Α	19870527		
os	MARPAT 110:38631				

- AB R1(R2A)NCH2CHR3CO2X [R1 = C1-22 (unsatd.) alkyl, alkoxyalkyl; R2 = C1-18 alkyl, C3-4 carboxyalkyl, carboxyphenyl, carboxy; R3 = H, Me; X = H, alkali metal, alkaline earth metal, (alkyl)ammonium alkanolammonium; A = C0, S02, CONH, C0-3 alkylene) useful as emulsifiers, wetteners, surfactants, and in preparation of waterproofing agents for leather, were prepared by reaction of R1NH2 with (meth)acrylic acid followed by acylation with carboxylic acid anhydrides, carbonyl chlorides, isocyanates, etc. Thus, CH2:CHCO2H was added to oleylamine at 60° and after 2.5 h and 90° maleic anhydride was added and the mixture was stirred for a further 2 h at 70-80° to give N-oleyl-N-(2-carboxyethyl)maleamic acid. The latter was used to prepare waterproof leather.
- L8 ANSWER 11 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 1987:554254 CAPLUS
- DN 107:154254
- TI Preparation and formulation of quinolonecarboxylic acid derivatives as antibacterials
- IN Enomoto, Hiroshi; Kise, Masahiro; Ozaki, Masakuni; Kitano, Masahiko; Morita, Iwao
- PA Nippon Shinyaku Co., Ltd., Japan
- SO U.S., 30 pp. CODEN: USXXAM
- DT Patent LA English
- FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI US 4659734	Α	19870421	US 1983-523329	19830815
CA 1259988	· A1	19890926	CA 1983-434945	19830819
PRAI US 1983-523329		19830815		

- OS MARPAT 107:154254
- GI For diagram(s), see printed CA Issue.
- The title compds. I [R1 = H, alkali metal, alkaline earth metal, alkyl, pivaloyloxymethyl, phthalidyl; R2 R5 = H, halo, alkoxy, disubstituted amino (including rings); A = (un)saturated hydrocarbyl; alkoxy, alkylthio, HO, halo; (un)substituted alkyl, O2N, NC, (un)substituted Ph, etc.) and their salts, were prepared Et 7-chloro-6-fluoro-4-hydroxy-2-mercaptoquinoline-3-carboxylate in DMF was refluxed with K2CO3 and BrCH2CH(OEt)2 and H2SO4 to give the Et thiazoloquinolinecarboxylate which was saponified to the free acid and substituted with N-methylpiperazine to give 7-fluoro-8-(4-methyl-1-piperazinyl)-5-oxo-5H-thiazolo[3,2-a]quinoline-4-carboxylic acid (II). II had an ED50 of 16 mg/kg against Pseudomonas aeruginosa in mice.
- L8 ANSWER 12 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 1987:176623 CAPLUS
- DN 106:176623
- TI Cyclopentadienylruthenium and -osmium complexes. IV. Separation and identification of ruthenocenecarboxylic and -boronic acids. Use and evaluation of the chromatographic test for the detection of complexing of alkali metal cations
- AU Wilczewski, Tadeusz
- CS Inst. Inorg. Chem. Technol., Tech. Univ. Gdansk, Gdansk, 80-952, Pol.
- SO Journal of Organometallic Chemistry (1986), 306(1), 125-32 CODEN: JORCAI; ISSN: 0022-328X
- DT Journal
- LA English
- OS CASREACT 106:176623
- AB Ruthenocene mono- and dicarboxylic acids were separated and identified. The applicability range of the chromatog. test, previously used to detect the complexing phenomenon of alkali metal cations by crown ethers, was determined The performance of the test in the case of several new cyclopentadienyl Ru and Os complexes, organic acids, and compds. of the ionic-pair type containing a large BPh4- anion, was investigated.
- L8 ANSWER 13 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 1984:209399 CAPLUS
- DN 100:209399

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2',4'-Difluoro-4-hydroxy-(1,1'-diphenyl)-3-carboxylic
ΤI
     Meneghin, Mariano; Piccinelli, Piero; Giordano, Claudio
IN
PA
     Zambon S.p.A., Italy
SO
     Eur. Pat. Appl., 6 pp.
     CODEN: EPXXDW
DT
     Patent
LA
     English
FAN.CNT 1
     PATENT NO.
                      KIND DATE
                                                                 DATE
                                         APPLICATION NO.
                                                                . -----
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                                          _____
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                               _____
    EP 101625 A1 19840229
EP 101625 B1 19860102
                               19840229 EP 1983-201056
ΡI
                                                                  19830716
       R: AT, BE, CH, DE, FR, GB, IT, LI, LU, NL, SE
               E 19860115 AT 1983-201056
2 A2 19840223 JP 1983-135051
7 B4 19930521
    AT 17230
                                                                  19830716
     JP 59033242
                                                                  19830722
     JP 05034347
    US 4486599
                        A
                               19841204
                                           US 1983-516356
                                                                  19830722
PRAI IT 1982-22516
    IT 1982-22516 A 19820722
EP 1983-201056 A 19830716
     The title acid, Diflunisal, was prepared from 4-(2,4-F2C6H3)C6H4OH (I) and
AB
     alkali metal alkyl carbonates; Diflunisal is useful as
     an antiinflammatory agent (no data). I was added to MeOC(O)ONa in MeOH,
     the MeOH was distilled, and the residue was heated under N at 200° to
     give Diflunisal.
L8
    ANSWER 14 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
AN
     1979:456370 CAPLUS
DN
     91:56370
ΤI
     1-Hydroperfluoroalken-(1)-yl carboxylic acid
     derivatives and their enol ethers and enol thioethers
IN
     Bathelt, Heinrich
PA
     Hoechst A.-G., Fed. Rep. Ger.
SO
     Ger. Offen., 20 pp.
     CODEN: GWXXBX
DT
     Patent
LA
     German
FAN.CNT 1
    PATENT NO. KIND DATE
                                         APPLICATION NO.
                                                                DATE
     _____
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                                          -----
                                                                  _____
PI DE 2742685 A1 19790405 DE 1977-2742685
PRAI DE 1977-2742685 A 19770922
                                                                  19770922
    RC(ZR1):CHCO2R3 [R = C1-11 perfluoroalkyl; R1 = alkyl, F3C(CF2)a(CH2)b (a
     is an integer of 0-16, b is an integer of 1-4), Ph, (CH2CH2O)nR2,
     (CH2CHMeO) nR2, (CHMeCH2O) nR2 (n = 1-50; R2 = H, alkyl); R3 = H, NH4+,
     alkali metal, alkyl; Z = O, S] and RCF:CHCO2R3 (R and
     R3 the same as above) were prepared Thus, dropping a 30% NaOMe solution into
     C8F17CH2CO2Me in MeOH over 20 min with ice cooling and stirring the mixture
     1 h at room temperature gave 94.2% C7F15C(OMe):CHCO2Me. Et3N was added to
     boiling C8F17CH2CO2Me in CCl2FCF2Cl over 10 min and the mixture was refluxed
     10 h to give 95% C7F15CF:CHCO2Me which was mixed with EtSH in CCl4.
     Treating this solution with Et3N at room temperature and refluxing the mixture 8 h
     gave 91% C7F15C(SEt):CHCO2Me.
L8
    ANSWER 15 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
ΑN
    1969:502264 CAPLUS
DN
    71:102264
     Chemical behavior of trialkylaluminum and dialkylzinc in the
TT
     copolymerization of \alpha-amino acid NCA[N-carboxylic
     acid anhydride] with epoxides
     Tsuruta, Teiji; Inoue, Shohei; Matsuura, Kazuo
ΑU
CS
     Univ. Tokyo, Tokyo, Japan
     Journal of Polymer Science, Polymer Symposia (1967), 22(Pt. 2), 981-92
SO
    CODEN: JPYCAQ; ISSN: 0360-8905
DT
    Journal
LA
    English
AΒ
    α-Amino acid-N- carboxylic acid anhydrides such
     as L(+)-alanine-NCA and L(-)-\beta-phenylalanine-NCA were copolymd. with
    propylene oxide by means of trialkylaluminum or dialkylzinc as catalyst.
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Structure of the copolymers formed were studied with the aid of ir and O.R.D. anal. Trialkylaluminum caused the formation of copolymers having more randomly distributed monomeric units of the amino acid. On the basis of results of the reaction modes of the two metal alkyls with the NCA group, the difference between the chemical behaviors of the metal alkyls in the copolymn. is discussed.

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ANSWER 16 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
1.8
AN
    1965:59047 CAPLUS
    62:59047
DN
OREF 62:10459f-h
    Organoboron mono and dicarboxylic acids and their preparation
    Ager, John W., Jr.; Alexander, Roy P.; Heying, Theodore L.
    Olin Mathieson Chemical Corp.
so
    6 pp.
DT
    Patent
LA
    Unavailable
FAN.CNT 1
                      KIND DATE APPLICATION NO.
    PATENT NO.
                                                                DATE
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                                          -----
                                                                 -----
PΙ
    US 3167584
                               19650126 US 1959-809569
                                                                 19590428
                              19590428
PRAI US
    The title compds. were prepared by treating RR1B10H8(CR2CR3) successively
    with an alkali metal alkyl or aryl, CO2, and an aqueous
    solution of a mineral acid. Thus, to a solution of BuLi (from 0.054 mole BuBr,
     2 hrs., at -10°) in 30 ml. Et2O was added a solution of 0.021 mole
    B10H10 (CHCH) in 20 ml. Et20 at 0°, the mixture brought to room temperature
     (precipitate formed), treated with CO2 (600 psi., 36 hrs.), extracted with 100 ml.
    H2O, and the aqueous solution acidified (HCl). The oil was extracted with Et2O and
    worked up to give a white pasty solid. Recrystn. from C6H6-heptane gave
     1.5 g. B10H10[C(CO2H)]2 m. 232°. Similarly prepared was
    B10H10[CH(CO2H)]2, m. 154-6°. The compds. were useful as fuels.
L8
    ANSWER 17 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
AN
    1955:60797 CAPLUS
DN
    49:60797
OREF 49:11717d-i,11718a-i
    Hydroxyhexahydrophenanthrenecarboxylic acids and derivatives
IN
    Hogg, John A.
PA
    Upjohn Co.
DT
    Patent
    Unavailable
LA
FAN.CNT 1
    PATENT NO.
                        KIND DATE
                                         APPLICATION NO.
                                                                 DATE
                             -----
                                          ______
                                                                 _----
PΙ
    US 2687426
                               19540824
                                         US 1947-756327
                                                                 19470621
GI For diagram(s), see printed CA Issue.
     I, where A is OH or a group convertible to OH by hydrolysis, B is CO2H or
     an esterified carboxyl group, and R is a lower alkyl, may be prepared by
     treating meta-substituted phenethyl halides with a Hagemann-type ester,
     reducing, cyclizing, and introducing the R group in the 2-position by
     treating with a metal alkyl. For example, 13 g.
     (0.563 mole) of Na was added portionwise to 250 ml. liquid NH3 containing 0.2
     g. hydrated Fe(NO3)3, cooling when necessary to speed addition The mixture was
     stirred until the blue color turned gray, the suspension was cooled in an
     alc.-Dry Ice bath, and 102.5 g. (0.563 mole) of Hagemann's. ester,
     1-methyl-6-carbethoxy-1-cyclohexen-3-one was added, with cooling.
     deep red mixture was stirred for 20 min. and cooled again while 300 ml. dry
     toluene and 50 ml. Na-dried ether were added. The mixture was stirred 2
     hrs. at room temperature until the NH3 had escaped and then heated to boiling,
     whereupon the yellow Na derivative precipitated m-Methoxyphenethyl bromide 120 g.
     (0.563 mole) was added and the suspension refluxed under N for 18 hrs.
     The mixture was washed with dilute HCl and H2O, the toluene layer was dried
     over MgSO4, and the toluene removed under vacuum. Distillation yielded 102 g.
     1-methyl-2-(m-methoxyphenethyl)-6-carbethoxy-1-cyclohexen-3-one (II), b.
     180-4°/0.3 mm. II 33 g. (0.104 mole) dissolved in 100 ml. of 95%
     EtOH, was hydrogenated in 45 min. under 35 lb. pressure, using 4 gm.
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palladized C as catalyst. A H2O-white oil of fruity odor was obtained

after filtration and removal of solvent. This oil was cooled to

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-20°, 80 ml. of cooled concentrated H2SO4 were added, the mixture was
stirred, and the temperature allowed to rise to below 10°. After 20 min.
of stirring, the temperature was allowed to rise to 20°, whereupon the
mixture was poured on excess ice. The total reaction time was 30 min. A
light colored gum, 1-methyl-2-carbethoxy-7-methoxy-1,2,3,4,9,10-
hexahydrophenanthrene, separated and was extracted with ether. The ether was
removed and the residue hydrolyzed by refluxing for 1 hr. in 200 ml. of 6%
KOH in 180 ml. of 95% EtOH and 20 ml. H2O. The alc. was vacuum distilled,
the residue diluted with H2O, washed with ether, and acidified with concentrated
HCl to precipitate the free acid as an oil which solidified; 2 recrystns. from
95% EtOH gave 14.2 g. 1-methyl-2-carboxy-7-methoxy-1,2,3,4,9,10-
hexahydrophenanthrene (III), m. 192-3°. When the preceding
procedure for preparing III from II was followed, omitting the hydrogenating
step, 1-methyl-2-carboxy-7-methoxy-3,4,9,10-tetrahydrophenanthrene (IV),
m. 192-5°, was obtained. The Me ester of IV, m. 112-13°C.,
was obtained by treating the acid with ethereal CH2N2. Demethylation of
II with HBr gave an oil, 1-methyl-2-carboxy-7-hydroxy-1,2,3,4,9,10-
hexahydrophenanthrene (V). III 7.5 g. was dissolved in ether, treated
with excess ethereal CH2N2, the solvent removed, and the residue recrystd.
from 95% EtOH to yield 7.3 g. Me 1-methyl-7-methoxy-1,2,3,4,9,10-
hexahydrophenanthrene-2-carboxylate (VI), m. 107-8°. Ph3CNa 0.0255
mole was added to an ether solution of 0.0255 mole VI in 300 ml. of Na-dried
ether, all under N. The wine-colored solution was allowed to stand at room
temperature with occasional shaking for 1 hr., the color fading to light orange.
Addition of 25 ml. MeI resulted in a reflux of the ether and precipitation of NaI.
After standing overnight, the ether was removed and the product,
1,2-dimethyl-2-carbomethoxy-7-methoxy-1,2,3,4,9,10-hexahydrophenanthrene,
was refluxed for 8 hrs. with 10 g. KOH in 10 ml. H2O and 200 ml. 95% EtOH.
The alc. was removed in vacuo the residue diluted with H2O to precipitate the K.
salt; the mixture of the salt and Ph3CH was filtered off and the latter
washed out with ether, leaving 7 g. white crystalline 1,2-dimethyl-2-carboxy-7-
methoxy-1,2,3,4,9,10-hexahydrophenanthrene (VII). The 1,2-diethyl- and
the 1-methyl-2-ethyl-2-carboxy-7-benzyloxy derivs. of VII were similarly
prepared from the corresponding derivs. of VI. Solution of the salt of VII in
250 ml. hot aqueous alc., acidification with HCl and recrystn. from 95% EtOH
gave 5.8 g. bundles of needles of the dl-trans form of the acid, m.
206-7°. HCl acidification of the aqueous alkaline filtrate from the preparation
of VII gave a tacky solid which was dissolved in 95% EtOH and allowed to
stand for 2 days to give the dl-cis form of the acid, m. 172-3° on
crystallization from EtOH. 1-Oxo-2-methyl-2-carbomethoxy-7-methoxy-1,2,3,4,9,10-
hexahydrophenanthrene (VIII) may be prepared by condensing
m-methoxyphenethyl halide and a suitably substituted \beta-oxopimelate,
cyclodehydrating the resulting ketone, hydrolyzing the resultant
dihydronaphthalene derivative, re-esterifying with an alc. and cyclizing with
a Na alcoholate to the Na salt of the unsatd. keto ester to give the
1-oxo-2-carbalkoxy-7-methoxyhexahydrophenanthrene. Treatment with Me
halide introduces the 2-methyl group to give VIII. Treatment of 12.5 g.
VIII in 200 ml. dry ether with 0.05 mole of EtMgBr in 50 ml. dry C6H6 at
room temperature for 3-4 min. gave the Me 1-hydroxy-1-ethyl-2-methyl- 7 -
methoxy - 1,2,3,4,9,10 - hexahydrophenanthrene - 2-carboxylate, which was
hydrolyzed with an excess of 1:1 concentrated HCl:H2O. The solvents were
removed, the residue solidified, and triturated with MeOH. Filtration and
crystallization from MeOH gave 8 g. of Me 1-ethylidene-2-methyl-7-methoxy-
1,2,3,4,9,10-hexahydrophenanthrene-2-carboxylate (IX), m. 145-7°.
Similar treatment of VIII with iso-PrMgBr gave the 1-isopropylidene derivative
of IX. VIII 8 g. in 200 ml. MeOH was reduced in an Adams hydrogenator
under 40 lb. pressure using 2 g. palladized C catalyst. After absorption
of I equivalent of H, the catalyst was removed and the product refluxed for 20
hrs. with a MeOH solution of 12 g. KOH dissolved in a min. of H2O.
solvent was removed, the product washed with cold water and filtered, and
dissolved in hot aqueous alc. Acidification gave an oil which solidified
while hot and melted 140-55°. Crystallization from EtOH gave crystals, m.
165-8°, and recrystn. gave 0.7 g. needles of 1-ethyl-2-methyl-2-
carboxy-7-methoxy-1,2,3,4,9,10-hexahydrophenanthrene (X), m.
175-6°. Hydrogenation of 200 mg. X at atmospheric pressure in 20 ml. MeOH
with 150 mg. palladized C catalyst until the theoretical amount of H was
absorbed, gave a white tacky gum after filtration and removal of solvent.
Trituration with 20 ml. of a 10% aqueous NaOH and filtration gave a filtrate A
and an insol. salt, which was dissolved in hot aqueous alc. and acidified to
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give 90 mg. 1-ethyl-2-methyl-2-carboxy-7-methoxy-1,2,3,4,9,10,11,12-octahydrophenanthrene (XI), m. 183-5°. Recrystn. from alc. gave chunky prisms, m. 188-90°. Acidification of filtrate A gave 100 mg. of acid, m. 60-65°. Trituration with 50% MeOH-H2O freed the residue of higher melting material, and on standing overnight at ice temperature deposited silky needles, a racemic-modification of XI, m. 112-15°, recrystd., m. 113-15°. These compds. are active estrogens in themselves as well as intermediates in the preparation of other estrogenic materials.

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ANSWER 18 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
L8
    1948:13793 CAPLUS
AN
     42:13793
DN
OREF 42:2995b-f
    Arylguanamines
TI
     Fairweather, Harold G. C.
IN
    American Cyanamid Co.
PA
DT
     Patent
     Unavailable
LA
FAN.CNT 1
                               DATE APPLICATION NO.
     PATENT NO.
                        KIND
                                                                 DATE
                               '-----
                                           _____
PΙ
    GB 593019
                                19471007
                                           GB
     Guanamines substituted in the 2-position with an aryl carboxylic
AB
     acid or sulfonated derivative thereof or an arylguanamine and with or
     without an alkenyl, aryl, or dialkyl substituent on the 6-NH2 group are
     prepared by treating a biguanide XYNC(:NH)NHC(:NH)NH2, where X is H,
     alkenyl, aryl, or Y when Y is alkyl and Y is H, or alkyl when X is also
     alkyl, with a dialkyl aryldicarboxylate or sulfonated derivative or with an
     alkali-metal alkyl aryldicarboxylate. The reaction is
     conducted in a lower (1-4 C atoms) aliphatic alc. with or without a metal
     alkoxide condensing agent. The products condensed with CH2O form resins
     soluble in dilute alkali, which, when applied to textiles and heat-cured, serve
     as water repellents. To 30 g. biguanide (I) in 100 g. MeOH is added 29 g.
     o-C6H4-(CO2Me)2. The mixture crystallizes on stirring and the biguanide
     salt of o-(4,6-diamino-s-triazin-2-yl)benzoic acid, m. 127-8° (from
     88% alc.), is filtered off in 75% yield; acidification of an aqueous solution to
     pH 4 with H2SO4 gives the free acid (II), m. 248-9°. Similarly, an
     equivalent amount of 2-ethylhexyl Me phthalate, b0.5 151-5° (prepared by
     distilling MeOH from o-C6H4(CO2Me)2and BuCHEtCH2OH with a trace of
     p-MeC6H4SO3H), gives 65% II and traces of 2,2' - o - phenylenebis(4,6 -
     diamino - s - triazine); p-C6H4(CO2C4H9)2 gives 45% 2,2!-p-
     phenylenebis (4,6-diamino-s-triazine), m. over 300°; and
     1,2-C10H6(CO2Me)2 gives a mixture of 1(and 2)-(4,6-diamino-s-triazin-2-yl)-
     2(and 1)-naphthoic acids. o-C6H4(CO2Me)CO2Na (60.6 g.), 25 g. I, and 6 g.
     Na in 300 g. MeOH give 94% of the crude Na salt of II which is converted
     to II in 75% yield. The Na salt of di-Am 4-sulfophthalate (20 g.) added
     to 6 g. I and 1.6 g. Na in 60 g. MeOH gives 90% of the di-Na salt of
     2-(4,6-diamino-s-triazin-2-yl)-4-sulfobenzoic acid. Cf. C.A. 41, 6769d;
     42, 2286g.
     ANSWER 19 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
L8
AN
     1930:34800 CAPLUS
DN
     24:34800
OREF 24:3713g-i,3714a
     Preparation of organic compounds by electrolytic methods. II. Electrolytic
TΙ
     reduction processes
ΑU
     Glasstone, S.
     Industrial Chemist and Chemical Manufacturer (1930), 6, 201-5
SO
     CODEN: ICCMAE; ISSN: 0367-7133
DT
     Journal
LA
     Unavailable
     Examples are given of reduction reactions illustrative of the principles
AB
     outlined in C. A. 24, 788. The electrolytic reduction of nitrobenzene in
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acid and alc. media and the effects of changes in conditions on the

products and the yields are given in detail. The reduction of other nitro compds. is dealt with, and the possibilities attendant upon the reduction of mixts. of organic substances is pointed out. The reduction of compds. containing the-CO group by the use of high overvoltage cathodes, e. g., Pb,

Hg, Zn, Cd and Tl is discussed. In general, the reduction of an aliphatic ketonic compound at a Pb cathode in acid solution yields a pinacone and a secondary alc. while in alkaline solution the secondary alc. is the main product. Reduction may proceed to the hydrocarbon, e. g., propane from acetone when a Cd cathode is used. Aliphatic aldehydes are generally reduced to the corresponding alc.. in acid solution at Pb or Hg cathodes. Aromatic ketones generally yield a secondary alc. as the main product in acid, neutral or alkaline solns. Aromatic aldehydes in alkaline solution yield usually hydroxybenzoin derivs. but in acid solution a primary alc. results. A carboxylic acid may be reduced electrolytically to the alc. or aldehyde, although a chemical reduction of this type is rarely possible. Fatty acids are not easily reduced, but aromatic acids can be converted to the corresponding alcs. The addition of H to a double bond and to the :C:Nlinkage is also dealt with. Possible applications of electroreduction processes to industrial purpose, e. g., the production of azo dyes and metal alkyl compds. (e. g., PbEt4), the reduction of soft fats to hard fats are pointed out as well as processes which are already in com. use, e. g., the reduction of indigo to indigo white.

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L8 ANSWER 20 OF 20 CAPLUS COPYRIGHT 2006 ACS on STN
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AN 1928:15145 CAPLUS

ON 22:15145

OREF 22:1769f-i,1770a-c

TI Presumable mechanism of polymerizations by alkali metals (preliminary communication)

AU Ziegler, K.; Bahr, K.

SO Ber. (1928), 61B, 253-63

DT Journal

LA Unavailable

GI For diagram(s), see printed CA Issue.

Z. and B. accidentally made an observation which is very probably of material significance in polymerizations produced by alkali metals: alkali metal alkyls can add to conjugated double bonds or double bonds adjacent to a C6H6 nucleus as Grignard reagents add to C:O compds. In an attempt to prepare PhCHKCHKPh (I) from (PhCH:)2 and PhCKMe2 there was obtained, instead of the expected violet-brown I, difficultly soluble in Et2O, an Et20-soluble, orange-yellow compound (II) converted by CO2 into a mixture of 2 (presumably diastereomeric dl-) α, β, γ -triphenyl- γ methylvaleric acids (III), whence II must be PhCMe2CHPhCHKPh. This addition of metallo-organic compds. to pure C:C double bonds seems to be a reaction of quite wide applicability. Thus far, PhCKMe2 has been found to add to styrene, Ph2C:CH2, anthracene and (CH2:CMe-)2 but not to cyclohexene. the resulting K compds. are colored, those obtained from styrene and Ph2C:CH2 probably have the structures PhCHKCH2CMe2Ph and Ph2CKCH2CMe2Ph and for that formed from (CH2:CMe)2 the structure CH2:CMeCKMeCH2CMe2Ph (resulting from 1,2-addition) is given the preference. Finally, the compound formed from anthracene is assigned the structure C6H4CH(CMe2Ph).C6H4.CHK. That these substances are addition products of the metallo-organic compound with the hydrocarbon is shown beyond doubt by the composition of the Ag salts of the CO2H acids obtained from them with CO2. Z. and B. do not wish to imply that addition will always occur at conjugated double bonds or double bonds adjacent to a C6H6 nucleus; perhaps other factors, not yet disclosed by the scant available exptl. data, play a decisive role not all organic K compds. are equally reactive; e. g., Ph3CK shows but slight tendency to Again, the metal alkyl may act merely as alkali donator, with resulting double decomposition; thus, phenanthrene with PhCKMe2 gives the same compound as was obtained by Schlenk with Na metal. The reaction readily explains how substances of very high mol. weight may be formed, for if a K alkyl, RK, reacts with, say, a butadiene, it forms again a K alkyl, RCH2KCH:CH2, with practically the same reactivity, which can add similarly to another butadiene mol. and the reaction can continue in this way until it is stopped by atmospheric O, traces of impurities or another butadiene mol. reacting according to the Shorigin scheme (RH + R'K = RK + R'H) or because the mols. finally become so large that they react only very sluggishly. The proposed scheme has been intentionally based on 1,2-addition to explain the difference between Na caoutchouc and natural caoutchouc; 1,4-addition would lead to products practically identical with natural rubber: In support of their views, Z. and B. have been able to isolate substances intermediate between the primary addition compound and the

final highly polymerized product. Below is the % of Ag in the salts of the CO2H acids obtained from the products of PhCKMe2 reacting with: 1 mol. styrene, 25.4; 3 mols., 19.3; 1 mol. (CH2:CMe)2, 29.36; 2 mols., 24.56; 1 mol. stilbene, 24.07; 1 mol. stilbene + 1 mol. styrene, 20.76. III, needles, m. 245-6°, and leaflets, m. 215-6°. α,α,γ -Triphenyl- γ -methylcaleric acid, from the addition product of PhCKMe2 to Ph2C:CH2, m. 154-6° (not quite pure; found for Ag salt, 23.01% Ag). 9-Phenylisopropyl-9,10-dihydroanthracene-10-carboxylic acid, from the addition product of PhCKMe2 to anthracene, m. 206-7°. Acid from the addition product of PhCKMe2 to phenanthrene, m. 223-4° (Ag salt, 32.92% Ag).

(FILE 'HOME' ENTERED AT 17:57:48 ON 08 MAY 2006)

FILE 'CAPLUS' ENTERED AT 18:00:43 ON 08 MAY 2006

=> s alkylation and photolysis 95706 ALKYLATION 103424 PHOTOLYSIS

L1 511 ALKYLATION AND PHOTOLYSIS

=> s l1 and carboxylic acid 237385 CARBOXYLIC 4142789 ACID 132188 CARBOXYLIC ACID

(CARBOXYLIC(W)ACID)

L2 8 L1 AND CARBOXYLIC ACID

=> d 1-8 bib abs

L2 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN

AN 2005:986132 CAPLUS

DN 143:440613

TI Synthesis of 17-epi-Calcitriol from a Common Androstane Derivative, Involving the Ring B Photochemical Opening and the Intermediate Triene Ozonolysis

AU Kurek-Tyrlik, Alicja; Michalak, Karol; Wicha, Jerzy

CS Institute of Organic Chemistry, Polish Academy of Sciences, Warsaw, 01-224, Pol.

SO Journal of Organic Chemistry (2005), 70(21), 8513-8521 CODEN: JOCEAH; ISSN: 0022-3263

PB American Chemical Society

DT Journal

LA English

OS CASREACT 143:440613

GΙ

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- AB An efficient synthesis of 17-epi-calcitriol (I), an epimer of natural hormone, via 17-epi-cholesterol II is described. Synthesis of II includes palladium-catalyzed cyclopropanation of the common androstane derivative III with an alkyl diazoacetate, reductive fission of the less shielded side of cyclopropane carboxylic acid esters, oxidation of the products into acid, and alkylation of ester. Photolysis of 7,8-dedydro-17-epi-25-hydroxycholesterol and consecutive thermal rearrangement gave a mixture of several products that was subjected to ozonolysis to provide, after chromatog., hydroxy ketone IV. The silyl derivative of IV was coupled with the resp. ring A building block.

RE.CNT 68 THERE ARE 68 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L2 ANSWER 2 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN

AN 2005:965868 CAPLUS

DN 143:413385

TI Sensitization of nanocrystalline TiO2 films with carboxy-functionalized bis(indoly1) maleimide

AU Kaletas, Basak Kuekrer; Kozhevnikov, Valery N.; Zimine, Mikhail; Williams, Rene M.; Koeniq, Burkhard; De Cola, Luisa

CS Molecular Photonic Materials, van't Hoff Institute for Molecular Sciences, Universiteit van Amsterdam, Amsterdam, 1018 WV, Neth.

SO European Journal of Organic Chemistry (2005), (16), 3443-3449 CODEN: EJOCFK; ISSN: 1434-193X

PB Wiley-VCH Verlag GmbH & Co. KGaA

DT Journal

LA English

AB The immobilization on a semiconductor surface of a bis(indolyl)maleimide

functionalized with two carboxylic acid groups by alkylation of the indole nitrogen atoms is presented and its synthesis is described. The compound, 3,4-bis[1-(carboxymethyl)-3-indolyl]-1H-pyrrole-2,5-dione is strongly colored and emissive and the imide part can coordinate an oxidizable substrate. Its absorption and emission spectra on TiO2 are substantially changed as compared to the spectra obtained in neat acetonitrile, indicating surface bonding through the carboxy groups. The quenching of the fluorescence of the sensitizer by the TiO2 surface is almost complete, reflecting the high degree of association between the TiO2 and the dye, fast charge injection and good electronic coupling between the sensitizer and the semiconductor. Nanosecond transient absorption spectra of the free sensitizer and of TiO2 surface bound sensitizer are recorded and compared. While the free chromophore in neat acetonitrile shows a transient absorption spectrum that decays on the nanosecond timescale (like the emission), the transient absorption spectra of the sensitized TiO2 film show a band at 360 nm, and a decay on the microsecond time scale. This is assigned to a slow recombination reaction of the charge-separated state. The properties discussed indicate that the authors system can be considered as a model compound for the development of photocatalysts immobilized on surfaces.

THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 32 ALL CITATIONS AVAILABLE IN THE RE FORMAT

- ANSWER 3 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN L2
- AN2001:854194 CAPLUS
- DN136:216402
- TISynthesis and studies on spectroscopic as well as electron donating properties of the two alkoxy benzo[b]thiophenes
- ΑU Misra, T.; Ganguly, T.; Kamila, S.; Basu, C.; De, A.
- CS Department of Spectroscopy, Indian Association for the Cultivation of Science, Calcutta, 700032, India
- SO Spectrochimica Acta, Part A: Molecular and Biomolecular Spectroscopy (2001), 57A(14), 2795-2808 CODEN: SAMCAS; ISSN: 1386-1425
- PB Elsevier Science B.V.
- DTJournal
- LA English
- os CASREACT 136:216402
- AΒ Synthesis, characterization, steady state and time resolved, using time correlated single photon counting as well as laser flash photolysis techniques, spectroscopic investigations were made for two alkoxy benzo[b]thiophene mols.: 5-methoxybenzo[b]thiophene (5MBT) and 5-(methoxymethoxy)benzo[b]thiophene (5MMBT). In both non-polar n-heptane (NH) and polar acetonitrile (ACN) solvents and at ambient temperature the electronic absorption spectra of these thiophenes exhibit different band systems whose assignments were made from the measurements of the steady state excitation polarization spectra. Steady state fluorescence spectra of these mols. in the different polarity solvents show the presence of non-specific interactions. From the redox properties of the benzothiophenes, measured by cyclic voltammetry, their electron donating properties were observed in the presence of the well-known electron acceptor 9-cyanoanthracene (9CNA). Further, detailed studies by laser flash photolysis techniques show that ion-recombination mechanism predominates after the initial excitation of the acceptor moiety using the third harmonic of Nd: YAG laser. This recombination together with the external heavy atom effect (the donor containing 'sulfur' atom) appears to be responsible for the formation of the triplet of the monomeric acceptor 9CNA. From the steady state expts. it is shown that both in non-polar NH and highly polar ACN the quenching in the fluorescence emission of 9CNA in the presence of the benzothiophene donors is brought about primarily by the external heavy atom effect and in ACN, although the presence of the photoinduced ET reaction is confirmed, this process seems, from the observed bimol. dynamic quenching rate, kq, to be significantly masked by the external heavy atom effect.
- THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 29 ALL CITATIONS AVAILABLE IN THE RE FORMAT
- ANSWER 4 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN 1.2 AN 2000:648011 CAPLUS

DN 133:296106

- TI Enantioselective photoelectrocyclization of a tropolone derivative in the crystalline state
- AU Scheffer, John R.; Wang, Letian
- CS Department of Chemistry, University of British Columbia, Vancouver, BC, V6T 1Z1, Can.
- SO Journal of Physical Organic Chemistry (2000), 13(9), 531-538 CODEN: JPOCEE; ISSN: 0894-3230
- PB John Wiley & Sons Ltd.
- DT Journal
- LA English
- Achiral tropolone ethers are well known to undergo photochem. induced ABdisrotatory electrocyclic ring closure in solution to form racemic bicyclo[3.2.0]hepta-3,6-dien-2-one derivs. The present paper reports successful efforts to carry out this transformation enantioselectively through the use of the solid-state ionic chiral auxiliary method. In this method, the reactant, an achiral tropolone ether, is equipped with a carboxylic acid group to which an optically pure amine can be attached by salt formation. Salt such as these are required to crystallize in chiral space groups, which provide asym. reaction cavities capable of differentiating enantiomeric transition states. Irradiation of these materials in the solid state leads to enantiomerically enriched products in moderate to high enantiomeric excess depending on the amine Of the amines studied, the best results were obtained with 1-phenylethylamine and 1-amino-2-indanol, which gave enantiomeric excesses in the 60-80% range depending on the extent of conversion. Because the tropolone ring is planar, it is suggested that the stereochem. outcome of the electrocyclization in the solid state is governed by environmental crystal lattice effects rather than by the initial conformation of the reactant.
- RE.CNT 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L2 ANSWER 5 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 1998:50046 CAPLUS
- DN 128:114560
- TI o-Nitrobenzyl as a photocleavable nitrogen protecting group for indoles, benzimidazole, and 6-chlorouracil
- AU Voelker, Troy; Ewell, Tim; Joo, Jean; Edstrom, Eric D.
- CS Department of Chemistry, University of Montana, Missoula, MT, 59812, USA
- SO Tetrahedron Letters (1998), 39(5/6), 359-362 CODEN: TELEAY; ISSN: 0040-4039
- PB Elsevier Science Ltd.
- DT Journal
- LA English
- OS CASREACT 128:114560
- AB The potential for the o-nitrobenzyl group as an alternative nitrogen protecting group for various indoles, benzimidazole, and 6-chlorouracil was determined Treatment of the appropriate N-H containing substrate with LiH or NaH in DMF followed by o-nitrobenzyl bromide afforded reasonable yields of N-alkylated products. To effect removal of this group, simple photolysis with 300 nm light afforded good yields of starting substrate.
- RE.CNT 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L2 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 1991:559666 CAPLUS
- DN 115:159666
- TI Mechanistic studies on DNA photolyase. 3. The trapping of the one-bond-cleaved intermediate from a photodimer radical cation model system
- AU Burdi, Doug; Begley, Tadhg P.
- CS Dep. Chem., Cornell Univ., Ithaca, NY, 14853, USA
- SO Journal of the American Chemical Society (1991), 113(20), 7768-70 CODEN: JACSAT; ISSN: 0002-7863
- DT Journal
- LA English

DNA photolyase catalyzes the monomerization of thymine photodimers in UV-damaged DNA. With the view to developing mechanistic probes for the enzymic reaction, the quinone sensitized cleavage of 6-iodomethyl photodimer model I system was examined This substituent proved to be an efficient trap for the one-bond-cleaved radical intermediate. The x-ray structural parameters for I were determined

L2 ANSWER 7 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1988:5813 CAPLUS

DN 108:5813

TI Synthesis of 2S,3aS,7aS- and 2S,3aR,7aR-perhydroindole-2carboxylic acid derivatives from L-aspartic acid

AU Barton, Derek H. R.; Guilhem, Jean; Herve, Yolande; Potier, Pierre; Thierry, Josiane

CS Inst. Chim. Subst. Nat., CNRS, Gif-sur-Yvette, 91190, Fr.

Tetrahedron Letters (1987), 28(13), 1413-16

CODEN: TELEAY; ISSN: 0040-4039

DT Journal

LA English

OS CASREACT 108:5813

GI

SO

AB Alkylation of aspartic acid derivative RNR1CH(CO2CMe3)CH2CO2R2 (I, R = R1 = H, R2 = CH2Ph) with 3-bromocyclohexene gave N-cyclohexenyl derivative I (R = 3-cyclohexenyl, R1 = H, R2 = CH2Ph) in 89% yield. Acetylation or p-bromobenzoylation, followed by basic hydrolysis, gave acids I (R = 3-cyclohexenyl; R1 = Ac, 4-BrC6H4CO; R2 = H), which were converted to N-hydroxy-2-thiopyridone derivs. I [R = 3-cyclohexenyl; R1 = Ac,4-BrC6H4CO; R2 = 1-(2-thiopyridonyl)] (II). Photolysis of II gave radical cyclization products III and IV (R1 = Ac, 4-BrC6H4CO; R3 = CMe3; R4 = 2-pyridylthio) as mixts. of diastereomers in 49-69% yields. Separation and desulfurization of III and IV with Raney Ni gave III and IV (R1 = Bz, R3 = CMe3, R4 = H). The stereochem. of III was determined by a crystal structure. Hydrolysis and re-esterification gave III and IV (R1 = Bz, R3 = Et, R4 = H). Separation desulfurization, and hydrolysis of III and IV (R1 = Ac, R3 = CMe3, R4 = 2-pyridylthio) gave amino acids III and IV (R1 = R3 =R4 = H), which were esterified and p-bromobenzoylated to give III and IV (R1 = 4-BrC6H4CO, R3 = Et, R4 = H).

- L2 ANSWER 8 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 1980:76898 CAPLUS
- DN 92:76898
- TI Carboxyl group-activated α-functionalized carboxylic

acid derivatives. I. A new route to methyl esters of N-pyruvoyl
amino acids
Burger, Klaus; Eggersdorfer, Manfred
Org. Chem. Inst., Tech. Univ. Muenchen, Garching, D-8046, Fed. Rep. Ger.
Liebigs Annalen der Chemie (1979), (10), 1547-53

CODEN: LACHDL; ISSN: 0170-2041
DT Journal
LA German

OS CASREACT 92:76898

GI

ΑU

CS SO

Oxazolidinones I (R = H, Me2CH, Ph, PhCH2) were dehydrogenated by photolysis in the presence of Br2 or by heating with SO2Cl2 to give the oxazolones II. II (R = Me, Et) were prepared by alkylation of II (R = H) with CH2N2 or CH3CHN2. II (R = Me) was treated with H-X-OMe (X = Gly, Ala, Val, Phe) to give 50-72% MeCOCO-X-OMe. II (R = Me, Et, Ph, PhCH2) were treated with H2NPh to give RCOCONHPh, and II (R = H, Me, Et, Me2CH, Ph, PhCH2) were cyclized with 1,2-(H2N)2C6H4 to give quinoxalines III. II are carbonyl group protected, carboxyl group activated, α-oxo carboxylic acid derivs.

=> s l1 and metal alkyl
1644131 METAL
567045 ALKYL
1739 METAL ALKYL
(METAL(W)ALKYL)
L3 3 L1 AND METAL ALKYL

=> d 1-3 bib abs

L3 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2006 ACS on STN

AN 1992:174407 CAPLUS

DN 116:174407

TI Carbene migratory insertions: preparation of the first second-row late-transition-metal alkyl-substituted carbene complex and comparison of its migratory insertion with that of its iron analog

AU Trace, Rhonda L.; Sanchez, Javier; Yang, Jing; Yin, Jianguo; Jones, W. M.

Dep. Chem., Univ. Florida, Gainesville, FL, 32611-2046, USA

SO Organometallics (1992), 11(4), 1440-2

CODEN: ORGND7; ISSN: 0276-7333

DT Journal

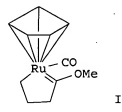
LA English

OS CASREACT 116:174407

GI

AB

CS



The first stable second-row late-transition-metal carbene complex substituted on the metal with an alkyl group, i.e., ruthenium complex I,

has been prepared Comparison with its iron analog shows carbene migratory insertion in the iron compound to be at least 107 times faster than in the ruthenium complex. This dramatic difference is ascribed to the stronger carbon-metal double bond in the ruthenium complex.

- L3 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 1987:576164 CAPLUS
- DN 107:176164
- TI Electronic absorption spectra and photochemical reactivity of Group 5 $metal\ alkyl$ compounds: photochemical α -hydrogen abstraction
- AU Chamberlain, Linda R.; Rothwell, Ian P.
- CS Dep. Chem., Purdue Univ., West Lafayette, IN, 47907, USA
- SO Journal of the Chemical Society, Dalton Transactions: Inorganic Chemistry (1972-1999) (1987), (1), 163-7 CODEN: JCDTBI; ISSN: 0300-9246
- DT Journal
- LA English
- OS CASREACT 107:176164
- The UV spectra and electrochem. and photochem. behavior of title compds. (RCH2)3MR12 [I; R = H, Me3Si, Ph; M = Nb, Ta; R1 = Cl, OCHMe2, OC6H3Me2-2,6, OC6H3(CHMe2)2-2,6, OC6H3(CMe3)2-2,6] were examined. The UV spectra of I are dominated by intense ligand-to-metal charge-transfer (LMCT) bands, the energy of which depends strongly on X. The photochem. of these compds., in some cases leading to the efficient and almost quant. photosynthesis of alkylidene functional groups, was examined. Irradiation of Me3Ta[OC6H3(CMe3)2-2,6]2 into the observed LMCT band at 313 nm produces the methylidene complex CH2:TaMe[OC6H3(CMe3)2-2,6]2 and CH4 with a quantum efficiency of 0.95 ± 0.1. Mechanistically the reaction is concerted, while for (Me3SiCH2)3Ta[OC6H3(CHMe2)2-2,6]2, photogeneration of the corresponding alkylidene involves an intermediate alkyl radical which can be intercepted.
- L3 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2006 ACS on STN
- AN 1987:554436 CAPLUS
- DN . 107:154436
- TI The synthesis and structure of Group 5 metal alkyl and alkylidene complexes containing 2,6-dialkylphenoxide ligands: x-ray crystal structures of [Ta(OC6H3Me2-2,6)2(CH2Ph)3], [Ta(OC6H3Me2-2,6)4Me], and [Ta(OC6H3Bu-tert2-2,6)2(:CHSiMe3)(CH2SiMe3)]
- AU Chamberlain, Linda R.; Rothwell, Ian P.; Folting, Kirsten; Huffman, John
- CS Dep. Chem., Purdue Univ., West Lafayette, IN, 47907, USA
- SO Journal of the Chemical Society, Dalton Transactions: Inorganic Chemistry (1972-1999) (1987), (1), 155-62 CODEN: JCDTBI; ISSN: 0300-9246
- DT Journal
- LA English
- OS CASREACT 107:154436
- AB Alkylation of Ta(OC6H3R2-2,6)3Cl2 (R = Me, CHMe2) with LiMe,
 LiCH2SiMe3 or Mg(CH2Ph)2 in C6H6 gave Ta(OC6H3R2-2,6)2(CH2R1)3 (I; R1 = H,
 SiMe3, Ph). Nb(OC6H3Me2-2,6)2Me3 was also prepared similarly.
 Photolysis of I (R = CHMe2, R1 = SiMe3) gave Ta[OC6H3(CHMe2)22,6]2(:CHSiMe3)(CH2SiMe3). Alkylation of Ta[OC6H3(CMe3)22,6]2Cl3 with LiCH2SiMe3 and Mg(CH2Ph)2 gave Ta[OC6H3(CMe3)22,6)2(:CHR1)(CH2R1) (II). The crystal structures of I (R = Me, R1 = H,
 Ph) and II (R1 = SiMe3) were determined